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(54) Coated metal electrodes for electrolysis

(57) An electrolytic electrode is made by coating an electrically conductive metal, e.g. Ti or an alloy thereof, with a solution of salts (e.g. chlorides) of Ti and/or Sn and of at least one of Al, Ga, Fe, Co, Ni and Tl, and heating the coated metal to oxidise the salts and form a corresponding mixed oxide coating; and then coating thereon an electrode-active substance, e.g. a Pt group metal oxide, e.g. by thermal decomposition.

The intermediate oxide layer should contain 5×10^{-3} mol/m² of metals; the ratio of (i) 4-valent Ti/Sn oxides to (ii) the other 2- or 3-valent oxide can be 95:5 to 60:40 by weight of metal; this layer presents passivation of the electrode and enhances adhesion of the electrode-active coating.

The electrode can be used as anode with a graphite plate as cathode, in an electrolyte, e.g. H₂SO₄, wherein O₂ is generated at the anode.

SPECIFICATION

Coated metal electrodes for electrolysis and process for production thereof

	Coated metal electrodes for electrolysis and process for production thereof	
5	The present invention relates to electrodes for electrolysis (hereinafter referred to as "electrolytic electrodes") and to a process for the production of the same. More particularly, the present invention relates to electrolytic electrodes having high durability, i.e., a long service life, when used in electrochemical processes, e.g., an aqueous solution in which the generation of oxygen at the anode is involved, and a process for the production of the same.	5
10	Heretofore, electrolytic electrodes comprising a substrate of valve metal, e.g., titanium (Ti), have been used as superior insoluble metal electrodes in the field of electrochemistry. In particular, they have been widely used as anodes for the generation of chlorine in the salt (sodium chloride) electrolytic industry. In addition to Ti, tantalum (Ta), niobium (Nb), zirconium (Zr), hafnium (Hf), vanadium (V), molybdenum (Mo), tungsten (W), etc. are known as valve	10
15	metals	15
	These metal electrodes are produced by coating metallic thanium with various electrochemically active substances such as platinum group metals and their oxides. Examples of such platinum group metals and their oxides are described in, e.g., U.S. Patent Nos. 3,632,498 and 3,711,385. These electrodes can maintain a low chlorine overvoltage over a long period of time as electrodes for the generation of chlorine.	20
	However, when the above metal electrodes are used as anodes in electrolysis for the generation of oxygen or electrolysis in which the generation of oxygen is involved, the anode overvoltage gradually increases. In extreme cases, the anode is passivated and thus it becomes impossible to continue the electrolysis.	25
25	The phenomenon of passivation of the anode is believed to be caused mainly by the formation of electrically non-conductive titanium oxides that result from (1) the oxidation of the titanium base material with oxygen by the electrode coating-consituting oxide substance itself; (2) oxygen diffusion-permeating through the electrode coating; or (3) the electrolyte. Formation of such electrically non-conductive oxides in the interface between the base	25
30	material and the electrode coating causes the electrode coating to peel off. This creates problems such as a breakdown of the electrode. Electrochemical processes in which the anode product is oxygen, or where oxygen is generated at the anode as a side reaction, include: (1) electrolysis using a sulfuric acid bath, a	30
35	nitric acid bath, an alkali bath or the like; (2) electrolytic separation of chromium (Cr), copper (Cu), zinc (Zn), or the like; (3) various types of electroplating; (4) electrolysis of dilute salt water, sea water, hydrochloric acid, or the like; and (5) electrolysis for the production of chlorate, and so forth. These processes are all industrially important. However, the above-described problems	35
40	have hindered metal electrodes from being used in these processes. U.S. Patent No. 3,775,284 discloses a technique to overcome passivation of the electrode due to permeation of oxygen. In this technique, a barrier layer of a platinum (Pt)-iridium (Ir) alloy, or of an oxide of cobalt (Co), manganese (Mn), lead (Pb), palladium (Pd), and Pt is provided between the electrically-conductive substrate and the electrode coating. The substances forming the intermediate barrier layer prevent the diffusion-permeation of	40
45	oxygen during electrolysis to some extent. However, these substances are electrochemically very active and therefore, react with the electrolyte passing through the electrode coating. This produces electrolytic products, e.g., gas, on the surface of the intermediate barrier layer which gives rise to additional problems. For example, the adhesion of the electrode coating is impaired due to physical and chemical influences of the electrode coating peeling off before the life of the substance of the electrode coating is over. Another problem is that the corrosion-resistance of	45
50	the resulting electrodes is poor. Thus, the method disclosed in U.S. Patent No. 3,775,284 fails to produce electrolytic electrodes which have high durability. Japanese Patent Application (OPI) No. 40381/76 (the term "OPI" used herein refers to a Published Unexamined Patent Application) discloses an intermediate coating layer comprising tin oxide doped with antimony oxide for coating the anode. However, the anode used is an anode	50
55	intended for the generation of chlorine, and hence an electrode provided with an intermediate coating forming substance disclosed in the above publication does not show the generation of oxygen.	55
60	U.S. Patent No. 3,773,555 discloses an electrode in which a layer of an oxide of, e.g., Ti, and a layer of a platinum group metal or an oxide thereof are laminated and coated on the electrode. However, this electrode has the problem that when it is used in electrolysis in which the generation of oxygen is involved, passivation occurs.	60
65	The present invention provides the ability to overcome the above-described problems. More specifically, an object of the present invention is to provide electrolytic electrodes which are especially suitable for use in electrolysis in which the generation of oxygen is involved, i.e., which resist passivation and have high durability.	65

	Another object of the present invention is to provide a process for producing such electrolytic electrodes.	
	According to the invention we provide:	
5	(I) an electrolytic electrode comprising (a) an electrode substrate of an electrically-conductive metal;	5
·	(b) an electrode coating of an electrode active substance; and	b
	(c) an intermediate layer provided between the electrode substrate (a) and the electrode	
	coating (b), wherein the intermediate layer (c) comprises a mixed oxide of	
	(i) an oxide of at least one member selected from titanium (Ti) and tin (Sn), each having a	
10	valence number of 4 in an amount of 60 to 95% by weight based on the weight of metal, and	10
	(ii) an oxide of at least one member selected from aluminum (AI), gallium (Ga), iron (Fe),	
	cobalt (Co), nickel (Ni), and thallium (TI), each having a valence number of 2 or 3 in an amount of 5 to 40% by weight based on the weight of metal; and	
	(II) a process for producing an electrolytic electrode, comprising the steps of:	
15	(1) coating an electrode substrate of an electrically conductive metal with a solution containing	15
	(i) salt(s) of Ti and/or Sn, and	, 5
	(ii) salt(s) of at least one metal selected from Al, Ga, Fe, Co, Ni and Tl, to provide a coated	
	substrate;	
20	(2) heating to an oxidizing atmosphere the electrode substrate coated with the solution in step	
20	(1), thereby forming on the electrode substrate an intermediate layer comprising a mixed oxide of	20
	(i) an oxide of at least one member selected from Ti and Sn in an amount of about 60 to	
	about 95% by weight based on the weight of metal, and	
	(ii) an oxide of at least one member selected from Al, Ga, Fe, Co, Ni and Tl, in an amount of	
25	about 5 to about 40% by weight based on the weight of metal; and	25
	(3) subsequently coating the intermediate layer with a layer of an electrode active substance.	
	The present invention is based on the discovery that the provision of the intermediate layer	
	between the substrate and the electrode coating enables one to obtain an electrode which can	
30	be used with sufficient durability as an anode for electrochemical processes in which the generation of oxygen is involved.	20
30	The intermediate layer in the present invention is corrosion-resistant and is electrochemically	30
	inactive. A function of the intermediate layer is to protect the electrode substrate, e.g., Ti, so as	
	to prevent passivation of the electrode without reducing its electrical conductivity. At the same	
	time, the intermediate layer acts to enhance the adhesion or bonding between the base material	
35	and the electrode coating.	35
	Accordingly, the present invention provides electrolytic electrodes which have sufficient	
	durability when used in electrolysis for the generation of oxygen or electrolysis in which oxygen is generated as a side reaction. Such processes have heretofore been considered difficult to	
	perform with conventional electrodes.	
40		40
	In the production of the electrode substrate of the present invention, corrosion-resistant.	. •
	electrically-conductive metals, e.g., Ti, Ta, Nb, and Zr, and their base alloys can be used.	
	Suitable examples are metallic Ti, and Ti-base alloys, e.g., Ti-Ta-Nb and Ti-Pd, which have	
45	heretofore been commonly used. The electrode base material can be in any suitable form such	
75	as in the form of a plate, a perforated plate, a rod, or a net-like member. The electrode substrate of the present invention may be of a type coated with a platinum	45
	group metal such as Pt or a valve metal such as Ta and Nb in order to increase corrosion	
	resistance or enhance the bonding between the substrate and the intermediate laver.	
	The intermediate layer is provided on the above-described electrode substrate and comprises a	
50	mixed oxide of an oxide of Ti and/or Sn having a valence number of 4 and an oxide of at least	50
	one member selected from the group consisting of Al, Ga, Fe, Co, Ni and Tl having a valence	
	number of 2 or 3. An electrody comprising on electrody substants of an electrical electrody substants of an electrical electrody.	
	An electrolytic electrode comprising an electrode substrate of an electrically conductive metal such as Ti and an electrode coating of a metal oxide, wherein an intermediate layer of a mixed	
55	oxide of an oxide of Ti and/or Sn and an oxide of Ta and/or Nb is provided between the	55
	substrate and the electrode coating is disclosed in U.S. Patents Nos. 4,471,006 and	J J
	4,484,999. This electrode is resistant to passivation and excels in durability. The intermediate	
	layer used in the electrode exhibits good conductivity as an N-type semiconductor. However,	
60	since the intermediate layer has limited carrier concentration, further improvement with respect	
οU	to conductivity was desired.	60
	Due to the concept of providing an intermediate layer possessing much higher conductivity than the intermediate layer of the electrode of these patents, the present invention has made it	
	possible to produce an electrode which eliminates the drawback suffered by the electrode of	
	these patents and offers still higher conductivity and durability.	
65	As the substance constituting the intermediate layer in this invention, a mixed oxide of an	65

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5	oxide of Ti and/or Sn and an oxide of at least one of Al, Ga, Fe, Co, Ni and Tl has been demonstrated to suit the purpose of this invention and provide an outstanding effect. The substance of the intermediate layer provides excellent resistance to corrosion, exhibits no electrochemical activity, and possesses ample conductivity. The term "oxide" or "mixed oxide" is meant to embrace solid solutions of metal oxides and metal oxides which are nonstoichiometric or have lattice defects. As used in this invention, the expression "TiO2", "SnO2", "Al2O3", "Ga2O3", "FeO", "FeO3", "CoO", "Co2O3", "NiO", "Tl2O3", etc. and the term "mixed oxide" embrace solid solutions of such metal oxides and those metal oxides nonstoichiometric or having lattice defects, for the sake of convenience.	5
10		10
15	Specifically, any of the mixed oxides TiO ₂ -Al ₂ O ₅ , TiO ₂ -Ga ₂ O ₃ , SnO ₂ -FeO, SnO ₂ -CoO, TiO ₂ -SnO ₂ -Co ₂ O ₃ , TiO ₂ -SnO ₂ -NiO, TiO ₂ -Al ₂ O ₃ -Tl ₂ O ₃ , SnO ₂ -Ga ₂ O ₃ -Fe ₂ O ₃ and TiO ₂ -SnO ₂ -Al ₂ O-Sa ₂ -Ga ₂ O ₃ can be used advantageously to achieve an ample effect. The proportions of the component oxides of the mixed oxide are not specifically defined and a wide range of proportions may be used. For protected retention of durability and conductivity of	15
20	the electrode, it is desirable for the ratio of the oxide of the tetravalent metal to the oxide of the divalent or trivalent metal to be in the range of about 95:5 to about 60:40 by the weight of metal. When the content of the oxide of the divalent or trivalent metal is not more than about 5% by weight substantially no improvement is observed as to the performance of the electrode, and the durability of the electrode decreases with not less than about 40% by weight of the	20
25	oxide of divalent or trivalent metal. The formation of the intermediate layer in the electrode can be advantageously effected by the thermal decomposition method which comprises the steps of applying a mixed solution containing chlorides or other salts of component metals destined to make up the aforementioned intermediate layer to the metal substrate and then heating the coated substrate in an	25
30	atmosphere of an oxidizing gas at temperatures of about 350° to 600°C thereby producing a mixed oxide. Other methods may be adopted if desired so long as the method is capable of forming a homogeneous, compact coating. By the afore-mentioned thermal decomposition method, Ti, Sn, Al, Ga, Fe, Co, Ni and Tl are readily converted into their corresponding oxides. The amount of the substance of the intermediate layer to be applied to the substrate preferably exceeds about 5 × 10 ⁻³ mol/m² calculated as metal. If the amount is less than about	30
35	5 × 10 ⁻³ mol/m ² mentioned above, the intermediate layer consequently formed does not provide sufficient effects. The thus-formed intermediate layer is then coated with an electrode active substance which is	35
40	electrochemically active to produce the desired product. Suitable examples of such electrode active substances are metals, metal oxides or mixtures thereof, which have superior electrochemical characteristics and durability. The type of the active substance can be determined appropriately depending on the electrolytic reaction in which the electrode is to be used. Active substances particularly suitable for the above-described electrolytic processes in which the generation of oxygen is involved include: platinum group metal oxides, and mixed oxides of platinum group metal oxides and valve metal oxides. Typical examples include: Ir oxide, Ir	40
45	oxide-Ru oxide, Ir oxide-Ti oxide, Ir oxide-Ta oxide, Ru oxide-Ti oxide, Ir oxide-Ru oxide-Ta oxide, and Ru oxide-Ir oxide-Ti oxide. The electrode coating can be formed in any suitable manner, e.g., by thermal decomposition,	45
50	electrochemical oxidation, or powder sintering. A particularly suitable technique is the thermal decomposition method a described in detail in U.S. Patents Nos. 3,711,385 and 3,632,498. The exact reason why the provision of the intermediate layer, i.e., the layer of the mixed oxide of 4-valent and 2- or 3-valent metals, between the metal electrode substrate and the electrode active coating produces the above-described results is not well understood. However, while not desiring to be bound the reason is believed as follows.	50
55	Crystallographically, it is confirmed that AI, Ga, Fe, Co, Ni and TI are in substantially a 6-coordination state and the ionic radii of these metals in a 6-coordination state vary within the range between the value by about 10% larger than and the value by about 10% smaller than that of Ti or Sn. This indicates that the mixed oxides of the metals form a layer of a uniform, dense solid solution or mixed oxide composed mainly of a rutile type crystal phase. Since such	55
60	an intermediate layer has a high resistance to corrosion, the surface of the substrate covered with the dense metal mixed oxide intermediate layer is protected from oxidation, and hence passivation of the substrate is prevented. In the intermediate layer, the 4-valent and 2-or 3-valent metals are present simultaneously as oxides. Therefore, according to generally known principles of Controlled Valency, the intermediate layer becomes an p-type semi-conductor having a very high electrical conductivity.	60
65	Moreover, where metallic Ti, for example, is used as a substrate, even when electrically non-conductive Ti oxides are formed on the surface of the substrate during the production of the	65

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below.

electrode or during the use of the electrode in electrolysis, the 2- or 3-valent metal in the intermediate layer diffuses and renders the Ti oxides semi-conductors. Accordingly, the electrical conductivity of the electrode is maintained and passivation is prevented. In addition, the intermediate layer substance which is composed mainly of rutile type oxides 5 enhances the adhesion or bonding between the substrate of, e.g., metallic Ti, and the electrode 5 active coating of, e.g., platinum group metal oxides and valve metal oxides, and hence increases the durability of the electrode. The present invention is described in greater detail by reference to the following examples which are in no way intended to limit the present invention. Unless otherwise indicated the 10 ratios are by weight. 10 **EXAMPLE 1** A commercially available Ti plate having a thickness of 1.5 mm and a size of 50 mm × 50 mm was degreased with acetone. Thereafter, the plate was subjected to an etching treatment 15 using a 20% aqueous hydrochloric acid solution maintained at 105°C. The thus treated Ti plate 15 was used as an electrode substrate. A 10% hydrochloric acid mixed solution of cobalt chloride, containing 10 g/l of Co, and titanium chloride containing 10.4 g/l of Ti, was coated on the Ti plate electrode substrate and dried. Thereafter, the plate was heated for 10 minutes in a muffle furnace maintained at 450°C. 20 This procedure was repeated five times to form an intermediate layer of a 4.0×10^{-2} mol/m² 20 TiO₂-Co₂O₃ mixed oxide (weight ratio of Ti to Co = 88:12) on the Ti substrate. A butanol solution of iridium chloride containing 50 g/l of Ir was coated on the above-formed intermediate layer and heated for 10 minutes in a muffle furnace maintained at 500°C. This procedure was repeated three times to produce an electrode with Ir oxide, containing 2.0 g/m² 25 of Ir, as an electrode active substance. 25 With the thus-produced electrode as an anode and a graphite plate as a cathode, accelerated electrolytic testing was performed in a 150 g/l sulfuric acid electrolyte at 60°C, and at a current density of 100 A/dm². The results demonstrated that this electrode could be used in a stable manner for 150 hours. For comparison, an electrode was produced in the same manner as above except that the 30 intermediate layer was not provided . This electrode was also tested in the same manner as above. The results demonstrated that this electrode was passivated in 20 hours and could no longer be used. Further, an electrode was produced in the same manner as above except that instead of 35 TiO2-Co2O3, a mixed oxide of SnO2 doped with antimony oxide in an amount of 20% by weight 35 calculated as Sb₂O₃ was used as the intermediate layer. When tested in the same manner as above, this electrode showed peeling off of the electrode active substance layer in 45 hours and could no longer be used. 40 EXAMPLE 2 40 An electrode was produced in the same manner as in Example 1 except that an intermediate layer of a TiO₂-Al₂O₃ mixed oxide (weight ratio of Ti to Al = 87.7:12.3) was provided. The thus-produced electrode was tested in the same manner as in Example 1. The results demonstrated that this electrode could be used for longer than 60 hours. 45 45 **EXAMPLE 3** A commercially available Ti plate having a thickness of 1.5 mm and a size of 50 mm imes 50mm was degreased with acetone. Thereafter, the plate was subjected to an etching treatment using oxalic acid maintained at 80°C for 12 hours. The thus-treated Ti plate was used as an 50 electrode substrate. 50 Various electrodes were produced by coating the electrode substrate with the intermediate layer shown in Table 1 below and an electrode active substance in the same manner as in Example 1. RuO2-IrO2 (weight ratio of Ru to Ir is 50:50) was used as the electrode active substance for each electrode. These electrodes were subjected to accelerated electrolytic testing 55 to determine their durability as an anode. The accelerated electrolysis was performed in an 55 aqueous 100 g/l sulfuric acid solution as the electrolyte at 40°C and at a current density of 200 A/dm² with a graphite plate as the cathode. The results obtained are shown in Table 1

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5	Run No.	Intermediate Layer	Service Life (hours)
	1	TiO ₂ -SnO ₂ -Fe ₂ O ₃ (22.8: 70.6: 6.6)	70
10	2	TiO ₂ -SnO ₂ -NiO (30.5:63.5:6.0)	64
10	3	$TiO_2-SnO_2-Ga_2O_3$ (25.3:47.1:27.6)	48
	4	$SnO_2-Co_2O_3$ (82.5:17.5)	54
15	5	SnO₂Tl₂O₃´ (70.0:30.0)	60
	6 (Comparison)	TiO ₂	30
20	7 (comparison)	$SnO_2 - Sb_2O_3$ (80: 20)	18

Note: The numerical values given in parentheses represent the weight ratios of component metals present.

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From the results in Table 1, it can be seen that electrodes incorporating an intermediate layer of this invention had decisively longer service life and exhibited higher durability than electrode (comparison) incorporating a conventional intermediate laeyr.

30 EXAMPLE 4

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Four electrodes as described in Table 2 below were produced in the same manner as in Example 1. These electrodes were subjected to accelerated electrolytic testing. The accelerated electrolytic tesing was perfored in a 12N aqueous NaOH solution at 95°C and at a current density of 250 A/dm². RuO₂-IrO₂ (weight ratio of Ru to Ir is 50:50) was used as the electrode 35 active substance for each electrode. The results are shown in Table 2.

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Table 2

40	Run No.	Intermediate Layer	Service Life (hours)
	1	TiO ₂ -SnO ₂ -Co ₂ O ₃ (10.4:76.9:12.7)	16
45	2	SnO ₂ -Fe ₂ O ₃ (90.6:9.4)	10
	3 (Comparison)		3
	4 (comparison)	$(SnO_2-Sb_2O_3) + SnO_2$ Powder (80: 20)	5

Note: The numerical values given in parentheses represent the weight ratios of component metals present.

It can be seen from the results in Table 2 that the electrodes of this invention have superior 55 durability and thus service life, to the comparative electrode.

As stated hereinabove, the electrodes of this invention shows excellent durability in electrochemical processes, particularly those in which generation of oxygen is involved, and can be used as various types of electrodes such as an electrolytic electrode and an electric cell or battery 60 electrode.

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CLAIMS

1. An electrolytic electrode comprising:

(a) an electrode substrate of an electrically-conductive metal;

(b) an electrode coating of an electrode active substance; and

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5	(c) an intermediate layer provided between the electrode substrate (a) and the electrode coating (b), wherein said intermediate layer (c) comprising a mixed oxide consisting of (i) an oxide of titanium and/or tin, each having a valence number of 4, in an amount of 60 to 95% by weight based on the weight of metal, and (ii) an oxide of at least one of aluminum, gallium, iron, cobalt, nickel and thallium, each having a valence number of 2 or 3, in an amount of 5 to 50% by weight based on the weight of metal.	5
	2. An electrode as claimed in Claim 1, wherein said electrode substrate (a) is one of	
10	titanium, tantalum, niobium or zirconium or an alloy thereof. 3. An electrode as claimed in Claim 1 or 2, wherein said intermediate layer (c) comprises an electroconductive mixed oxide of	10
	(i) TiO ₂ and/or SnO ₂ and	
	(ii) at least one oxide selected from Al ₂ O ₃ , Ga ₂ O ₃ , FeO, Fe ₂ O ₃ , CoO, Co ₂ O ₃ , NiO and Tl ₂ O ₃ . 4. An electrode as claimed in Claim 1, 2 or 3, wherein said electrode active substance	
15	contains a platinum-group metal or an oxide thereof. 5. An electrolytic electrode as claimed in Claim 1, substantially as hereinbefore described in	15
	any of Examples 1 to 4 apart from the comparison runs.	
	6. A process for producing an electrolytic electrode, comprising the steps of:	
20	(1) coating an electrode substrate of an electrically conductive metal with a solution containing (i) salt(s) of Ti and/or Sn, and (ii) salt(s) of at least one metal selected from Al, Ga, Fe, Co, Ni and Tl to provide a coated electrode substrate;	20
	(2) heating in an oxidizing atmosphere the electrode substrate coated with said solution in	
	step (1), thereby forming on said substrate an intermediate layer (c) as defined in Claim 1 or 3;	
25	(3) subsequently coating said intermediate layer with a layer of an electrode active substance.	25
	/. A process as claimed in Claim 6, wherein said coating of the intermediate layer with said	25
	electrode active substance is carried out by thermal decomposition.	
	8. A process as claimed in Claim 7, wherein said intermediate layer is formed by heating the coated electrode substrate under an oxidizing atmosphere at a temperature of 350° to 600°C.	
30	9. A process as claimed in Claim 6, 7 or 8, wherein said electrode substrate is one of	30
	titanium, tantalum, niobium, or zirconium or an alloy thereof. 10. A process as claimed in Claim 6, 7, 8 or 9, wherein said electrode active substance	
	contains a platinum-group metal or an oxide thereof.	
35	11. A process as claimed in Claim 6, for producing an electrolytic electrode, substantially as	
33	hereinbefore described in any of Examples 1 to 4 apart from the comparison runs. 12. An electrolysis process wherein an electrode as described in any of Claims 1 to 5 or	35
	made by a process as claimed in any of Claims 6 to 11 is used.	